## In the Claims

Please amend claims 1-14 as follows:

- 1. (Currently amended) A method of enhancement of an immune response and immunomodulating activity comprising administration to a subject an effective amount of an [[An]] adjuvant composition with synergistic effect[[,]] comprising[[:]] an iscom particle comprising
- [[a]] fraction  $\underline{A}$  of Quil A[[; and]] together with at least one other adjuvant[[,]] in free form or integrated into another separate iscom particle

wherein the composition provides enhanced immune responses and immunomodulating activity.

- 2. (Currently amended) The method An adjuvant composition according to claim 1 wherein said at least one other adjuvant is chosen from the group consisting of: saponins, naturally occurring saponin molecules derived from crude saponin extract of Quillaja saponaria Molina, synthetic saponin molecules derived from crude saponin extract of Quillaja saponaria Molina, semisynthetic saponin molecules derived from crude saponin extract of Quillaja saponaria Molin, saponin fractions from Quil A, saponin fractions from cell wall skeleton, blockpolymers, hydrophilic block copolymers, CRL-1005, Threhalose di mucolate (TDM), lipopeptides, LPS derivatives, LPS-derivatives, Lipid A from a bacterial species and derivatives thereof, monophosphoryl lipid A, CpG variants, CpGODN variants, endogenous human animal immunomodulators, GM-CSF. IL-2, native adjuvant active bacterial toxins, modified adjuvant active bacterial toxins, cholera toxin CT, CT subcomponent CTB, CT subcomponent CTA1, thermolabile toxin (LT) of E. coli, Bordetella pertussis (BP) toxin, and a filamentus heamagglutenin of BP.
- 3. (Currently amended) <u>The method An adjuvant composition</u> according to claim 2 wherein the saponin fraction from Quil A is fraction C of Quil A or fraction B of Quil A.

4. (Currently amended) The method An adjuvant composition according to claim 1, wherein said at least one other adjuvant is integrated into one iscom particle.

- 5. (Currently amended) The method An adjuvant composition according to claim 1, wherein said fraction A of Quil A is integrated into a first iscom particle and said at least one other adjuvant is integrated into a second iscom particle.
- 6. (Currently amended) The method An adjuvant composition according to claim 5, wherein said at least one other adjuvant is integrated into a plurality of separate iscom particles.
- 7. (Currently amended) The method An adjuvant composition according to claim 4, wherein said fraction A of Quil A is integrated into one iscom particle and said at least one other adjuvant is not integrated into iscom particle.
- 8. (Currently amended) The method An adjuvant composition according to claim 7, wherein said at least one other adjuvant is at least one of monophosphoryl lipid A and cholera toxin CT.
- 9. (Currently amended) The method An adjuvant composition according to claim 4, wherein said iscom particle is an iscom complex.
- 10. (Currently amended) The method An adjuvant composition according to claim 4, wherein said iscom particle is an iscom matrix complex.
- 11. (Currently amended) <u>The method</u> An adjuvant composition according to claim 3, wherein the composition comprises

50-99.9% of fragment A of Quil A; and

0.1-50% of a fraction or derivative of Quil A based on the total weight of the composition.

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wherein the composition comprises

- 12. (Currently amended) The method An adjuvant composition according to claim 11,
  - 75-99.9% of fragment A of Quil A; and
- 0.1-25% of a fraction or derivative of Quil A based on the total weight of the composition.
- 13. (Currently amended) The method An adjuvant composition according to claim 12, wherein the composition comprises
  - 91-99.1 % of fragment A of Quil A; and
  - 0.1-9% of a fraction or derivative of Quil A based on the total weight of the composition.
- 14. (Currently amended) The method An adjuvant composition according to claim 1, wherein the composition further comprises a pharmaceutically acceptable carrier, diluent, excipient or additive.